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2 CHAPTER TWO: MATERIALS

A HMA pavement is composed of binder and aggregate blended together. The individual material properties of each component can affect the overall performance of the pavement. If pavements are to perform long term and withstand specific traffic and loading, the materials making up the pavements must be of high quality.

This section will cover in detail what material properties are necessary for high quality pavements. The materials covered in this section are aggregates, and PG binders. Many of the tests are not required to be performed by the certified technician; however a thorough understanding of the materials and test procedures to determine quality is a necessary background for the certified technician. Converting that understanding into a working knowledge will assist the technician in making accurate, reliable day to day decisions.

AGGREGATES

Source

Aggregates for HMA are generally classified according to their occurrence. The three major sources of aggregates for INDOT use are sand and gravel operations, stone quarries, and industrial processing (synthetic aggregates). Crushed stone, sand, and gravel are classified as natural aggregates. Blast furnace slag, steel slag, granulated blast furnace slag, and wet bottom boiler slag are by-products of industrial processing and are classified as synthetic aggregates.

Sand and gravel generally occur together, where they were deposited by a stream or glacier. These aggregates may be produced either from a water-filled pit using a suction dredge or a crane and dragline bucket, or from a cut-back deposit, using end-loaders or shovels. The raw materials produced in this manner are, therefore, called either pit-run or bank-run materials. These materials require further processing, such as screening, washing and crushing, to produce aggregates of the proper size.

Crushed stone is produced from quarries where bedrock is blasted from a quarry face and then further processed by crushing and screening. Crushing is accomplished by processing the shot rock through a series of crushers until the desired top size is produced. Materials are then graded in a screening operation similar to the sand and gravel operations.

Synthetic aggregates are produced either from blast furnace slag, steel slag, or from wet bottom boiler slag. Blast furnace or steel slag is produced as a by-product of iron or steel production. These materials are non-metallic substances that rise to the surface of molten iron or steel during the smelting process. After being drawn off the surface of the melt, the slag is placed into a pit and is allowed to solidify by the prevailing atmospheric conditions. Granulated slag is blast furnace slag that has been solidified by quickly quenching the material in water. Wet bottom boiler slag is a by-product of coal-fired electric power plants and is commonly known as cinders.

Physical Qualities

Physical quality requirements for aggregates used in HMA are all specification provisions other than those dealing with gradation or usage requirements. These quality requirements can be divided into five distinct groups as follows:

- 1) Absorption/Specific Gravity;
- 2) Abrasion Resistance;
- 3) Soundness;
- 4) Deleterious Materials; and
- 5) Special Requirements

Absorption and Specific Gravity

The internal pore characteristics are very important properties of aggregates. The size, the number, and the continuity of the pores through an aggregate particle can affect the strength of the aggregate, abrasion resistance, surface texture, specific gravity, bonding capabilities, and resistance to freezing and thawing action. Absorption relates to the particle's ability to take in a liquid.

Density is the weight per unit of volume of a substance. Specific gravity is the ratio of the density of the substance to the density of water. The following chart illustrates these relationships for some common substances.

Substance	Specific Gravity	Density (lb/ft ³)
Water	1.0	62.4 lb/ft ³
Binder	1.02	63.7 lb/ft ³
Limestone	2.7	165 to 170 lb/ft ³
Lead	11.0	680 to 690 lb/ft ³

The density and the specific gravity of an aggregate particle is dependent upon the density and specific gravity of the minerals making up the particle and upon the porosity of the particle. These may be defined as follows:

- 1) All of the pore space (bulk density or specific gravity);
- 2) Some of the pore space (effective density or specific gravity); or
- 3) None of the pore space (apparent density or specific gravity).

Determining the porosity of aggregate is often necessary, but it is difficult to directly measure the volume of pore space. Correlations can be made between porosity and the bulk, apparent and effective specific gravities of the aggregate.

As an example, specific gravity information about a particular aggregate will help in determining the amount of binder needed in the HMA. If an aggregate is highly absorptive, the aggregate will continue to absorb binder after initial mixing at the plant, until the mix cools down completely. This process will leave less binder for bonding purposes; therefore, a more porous aggregate requires more binder than a less porous aggregate. The porosity of the aggregate can be taken into consideration in determining the amount of binder required by applying the three types of specific gravity measurements.

In the example in Figure 2-1 the bulk specific gravity includes all the pores, the apparent specific gravity does not include any of the pores that would fill with water during a soaking, and the effective specific gravity excludes only those pores that would absorb binder.

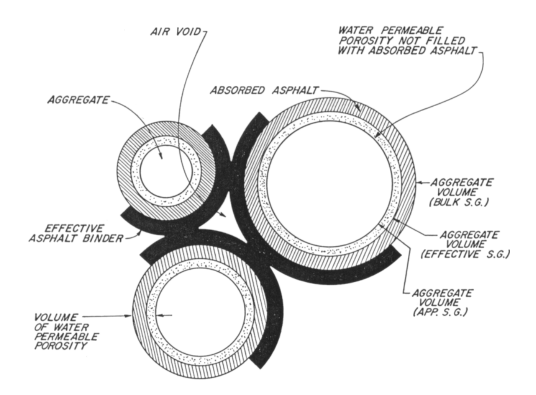


Figure 2-1. Aggregate Specific Gravities

Abrasion Resistance

For a coarse aggregate to perform satisfactorily in a pavement it must be tough enough to withstand the action of rolling during construction and the action of traffic without breaking down under the imposed loads. The test used for evaluating this property is the Los Angeles abrasion test (AASHTO T 96). Briefly, the aggregate is placed in a metal drum along with a charge of steel balls, and the drum is rotated 500 times. The inside of the drum is equipped with an angle iron which runs longitudinally. This causes the charge of aggregate and balls to fall with a heavy impact once during each revolution, breaking the aggregate particles into smaller particles. At the completion of the test the aggregate is shaken over a No. 12 sieve and the amount which passes through the sieve, expressed as a percentage of the total charge, is the Los Angeles abrasion value designated "percent wear". Very hard aggregates will have values of 20 percent or less; softer aggregates, such as a limestone which may be scratched easily with a pocketknife, may have values over 40 percent. Aggregates, with the exception of Blast Furnace Slag, with a wear of over 40 percent are unsatisfactory for use in surface HMA. Aggregate used in a base HMA may be softer than used in a surface HMA. The test equipment is illustrated in Figure 2-2.

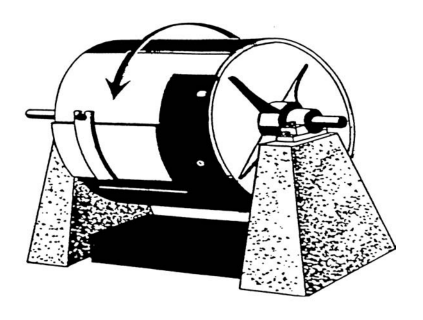


Figure 2-2. Los Angeles Abrasion Apparatus

Soundness

Aggregates which disintegrate badly under the force of weathering are termed unsound. Shale is a typical unsound material because water enters into it and freezes, causing it to expand and disintegrate. Also, exposure to air (oxidation) will cause shale to flake. Unsound aggregates are obviously unsatisfactory for use in HMA, particularly for surface HMA, which are more exposed to the weather. INDOT subjects aggregates to three different test methods to evaluate soundness:

- 1) The sodium sulfate test in test method **AASHTO T 104**;
- 2) The brine freezing and thawing test in **ITM 209**; and
- The freezing and thawing test in test method **AASHTO** T **103**.

The sodium sulfate test requires immersing an aggregate sample in a sodium sulfate solution for a period of time and then determining the weight loss of particles on a given set of sieves. The brine freeze and thaw test requires the aggregate to be enclosed in a bag containing a 3 percent sodium chloride solution and subjected to 25 cycles of freeze and thaw. The water freeze and thaw test requires the aggregate to be sealed and totally immersed in water and subjected to 50 cycles of freeze and thaw.

The freezing and thawing in water test is the method that most accurately simulates actual field conditions, but the test requires a long period of time to conduct. The "quick" checks for soundness of the aggregate are the brine freeze and thaw and sodium sulfate test. If the aggregate fails either the brine freeze and thaw or the sodium sulfate test, the material is tested using the freeze and thaw method with water. An aggregate that reasonably fails the brine freeze and thaw and/or sodium sulfate test but then passes the freeze and thaw in water method is an acceptable material for use on INDOT contracts.

Deleterious Materials

Certain substances in aggregates are undesirable for use in HMA. Therefore, the Standard Specifications limit the amount of deleterious constituents to a level consistent with the quality sought in the final products. Figure 2-3 illustrates the materials which are classified as deleterious and the specification limits for each. Of particular interest for avoidance in HMA is chert.

Chert is mineral matter composed of microcrystalline silica. When lightweight chert (less than 2.45 specific gravity) is present in aggregate used to produce HMA, the chert may undergo volumetric expansion sufficient to cause disruption of the mix, causing popouts, when the chert particles are located close to the surface. In addition, the binder is unable to coat the particles which leads to raveling of the pavement under traffic. Once either problem is initiated, the rate of deterioration of the surface is accelerated.

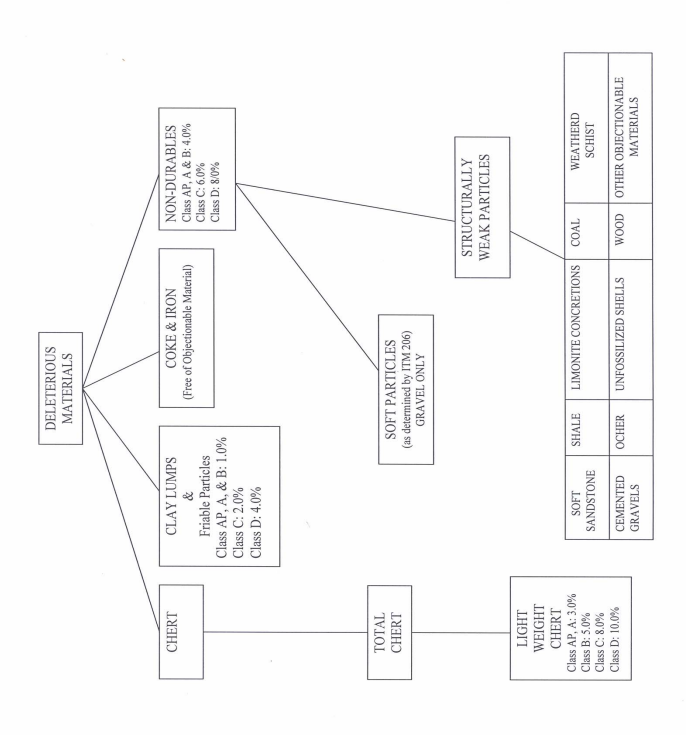


Figure 2-3. Deleterious Materials

Special Requirements

Particle Shape

The shape of the aggregate particles affect such things as:

- 1) The binder demands of HMA; and
- 2) The workability and the strength of HMA.

The best aggregates to use for strength are crushed stone or crushed gravel. Crushed aggregates have irregular, angular particles that tend to interlock when compacted or consolidated.

The crushed stone or crushed gravel aggregate make the HMA somewhat difficult to place. To improve the workability, many mixes contain both angular and round particles. The coarse aggregate particles are usually crushed stone or crushed gravel, and the fine aggregate particles are usually natural sand.

Surface Texture

Like particle shape, the surface texture also influences the workability and strength of HMA. Surface texture has often been considered more important than the shape of the aggregate particles. A rough, sandpaper-like surface texture as opposed to a smooth surface tends to increase the strength of the HMA. It is important to understand that some aggregates may initially have good surface texture but under traffic may polish smooth.

Coatings

Coating is a layer of substance covering a part or all of the surface of an aggregate particle. The coating may be of natural origin, such as mineral deposits formed in sand and gravel by ground water, or it may be artificial, such as dust formed by crushing and handling.

Generally, aggregates used in HMA must be washed to remove the coating (contaminant) left on the particles. The coating may prevent a good bond from forming between the aggregate surfaces and the binder. The coating could even increase the quantity of binder needed in the HMA. Also, if the quantity of the coating varies from batch to batch, undesirable fluctuations in the consistency of the HMA may result.

Coarse Aggregate Angularity

Crushed particles (coarse aggregate angularity) are necessary in HMA to assist in resisting shoving and rutting under traffic. The internal friction among the crushed aggregate particles prevents them from being moved past each other and provides for a stable mix. The test procedure used to define coarse aggregate angularity is **ASTM D 5821**.

The crushed particle requirement applies for HMA when gravel is used. Crushed particles are defined as those particles having one or more sharp, or slightly blunt edges. Fractured faces that have an area less than 25 percent of the maximum cross sectional area of the particle are not considered crushed.

Flat and Elongated Particles

Flat and elongated particles are undesirable because they have a tendency to break during construction and under traffic. This characteristic is defined as the percentage by weight of coarse aggregates that have a length in excess of five times its width in accordance with **ASTM D 4791**. Figure 2-4 is an illustration of the device used to measure these particles.

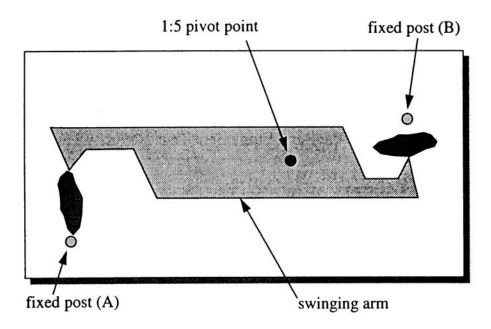


Figure 2-4. Flat and Elongated

Clay Content

Limitation of the amount of clay in aggregate strengthens the adhesive bond between the binder and the aggregate. Clay content is the percent of clay material contained in the aggregate fraction that is finer than a No. 4 sieve.

The test used for determining the clay content is the Sand Equivalent Test (AASHTO T 176). In this test, a sample of fine aggregate is placed in a graduated cylinder with a flocculating solution and agitated to loosen clayey fines present in and coating the aggregate. The flocculating solution forces the clayey material into suspension above the granular aggregate. After a period that allows sedimentation, the cylinder height of suspended clay and sedimented sand is measured (Figure 2-5). The sand equivalent value is computed as a ratio of the sand to clay height readings expressed as a percentage.

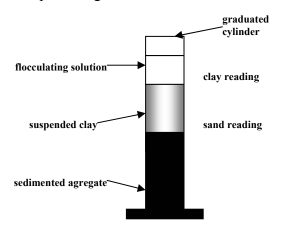


Figure 2-5. Sand Equivalent

Fine Aggregate Angularity

Fine aggregate angularity, like the crushed content of coarse aggregate, is necessary to achieve a high degree of internal friction and thus, high shear strength for rutting resistance. It is defined as the percent air voids present in loosely compacted aggregates finer than the No. 8 sieve.

The test used for determining the Fine Aggregate Angularity is the Uncompacted Void Content of Fine Aggregate Test (AASHTO T 304). In the test, a sample of fine aggregate is poured into a small calibrated cylinder by flowing through a standard funnel (Figure 2-6). By determining the weight of fine aggregate (W) in the filled cylinder of known volume (V), void content can be calculated as the difference between the cylinder volume and fine aggregate volume collected in the cylinder. The fine aggregate bulk specific gravity (Gsb) is used to compute fine aggregate volume.

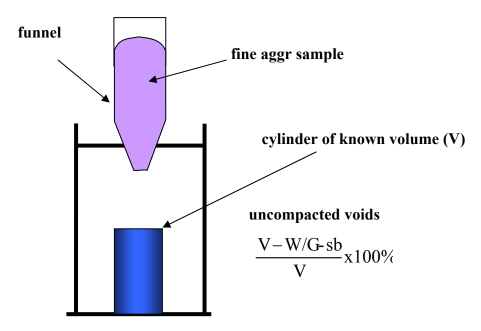


Figure 2-6. Fine Aggregate Angularity

Dolomitic Aggregates

There is a special requirement to be met when dolomitic coarse aggregates are used in HMA. These aggregates are specified under some conditions to obtain high-friction, skid-resistant HMA surface courses. **ITM 205** is used to ensure that the aggregate is a carbonate rock containing at least 10.3 percent elemental magnesium.

Polish Resistant Aggregates

Aggregates that meet the requirements of **ITM 214** may be used in place of dolomitic aggregates in HMA surface mixtures. The procedure for approval requires initial British Pendulum testing, placement of a test section on an INDOT project, and subsequent skid testing for two years.

Sandstone Aggregates

Coarse Sandstone must meet the Class B quality requirements, and may only be used in HMA surface mixtures. The definition of sandstone is described in Section **904.01** of the Standard Specifications.

Slag Aggregates

When slag is furnished as an alternate to natural aggregate, payment is made on a weight basis. Adjustments must be made to compensate for the difference in specific gravity of the slag compared to the specific gravity of the natural aggregate. For any pay item less than 500 tons on a contract, no adjustment is made. The following typical values should be used

TYPICAL VALUES FOR SPECIFIC GRAVITY				
Natural aggregates (both fine and coarse)	2.6			
Air cooled blast furnace slag coarse aggregate	2.3			
Air cooled blast furnace slag fine aggregate	2.6			
Granulated blast furnace slag fine aggregate	2.1			
Steel furnace slag, both fine and coarse	3.2			

Stockpiling

Segregation is probably the greatest problem of stockpiling and handling, but certainly other problems such as degradation and contamination can adversely affect product quality. Every possible precaution should be taken to protect product quality from initial stockpiling to the point where the material is loaded into the HMA plant.

The majority of aggregate stockpiles at the HMA plant are built by dumping individual truckloads of material. The best truck-built stockpiles are those that are constructed one dump high with each dump placed against previously dumped material. Here, because of the low profile, roll-down segregation is minimal and can be reduced by reasonable remixing effort during loading of the cold bins. However, these stockpiles require a large area. A technique that can help reduce the required area is to restock some dumps on top of other dumps with a large endloader operating from ground level. In this case, care should be taken to place the upper lift back from the edge far enough that a long sloped face is not created that would cause segregation.

Occasionally aggregate is dumped over a quarry or pit face to form a stockpile. This tends to cause considerable segregation, particularly with larger and long graded aggregates. In general, the larger particles work to the outside and base of the pile. The extent of segregation varies with the height of fall, gradation of the material, moisture, and other conditions. Segregation typically occurs as shown in Figure 2-7.

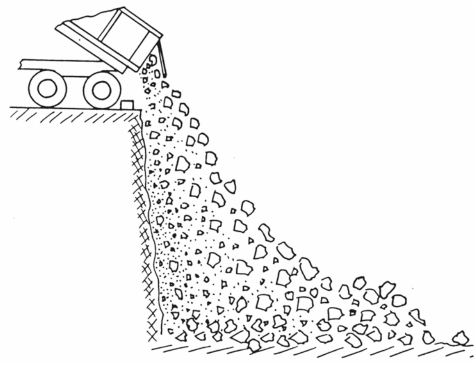


Figure 2-7

Sampling

Because of the various sampling locations and the availability of equipment, there are several methods of taking aggregate samples. Uniformity of obtaining the sample cannot be emphasized enough. The uniformity eliminates one variable in test results.

Bin Sampling

Bin samples taken at HMA plants should be taken at the discharge chute of the bin. In this case a number of small samples are taken at short intervals and combined to make the total sample. Each of these samples should include the entire cross section of the flow of material from the chute at the moment taken. If the rate of flow from the discharge chute cannot be controlled, it may be impractical or impossible to obtain this type of sample by passing a bucket across the flow of material. Therefore, a mechanical sampling device that will support the weight of sample or a diversion chute may be required.

Belt Sampling

Belt sampling material consist of taking samples of materials directly from conveyor belts. The proper procedure is to:

- 1) Make sure that the belt is carrying a normal load of material which is not segregated;
- 2) Have the plant operator stop the belt;
- Take a complete cross section of the material, being careful to include all the material on the belt and only the material in the section. A template is recommended. Remove most of the sample with a scoop or shovel and the remainder with a brush; and
- 4) Take as many complete cross sections as necessary to obtain a sample that meets the minimum sample size.

Stockpile Sampling

The primary control of aggregates at the HMA plant is accomplished by sampling stockpiles of the finished materials. Therefore, it is important to establish a uniform and representative method of sampling (ITM 207). Coarse aggregate samples are taken with a large square nose shovel; fine aggregate samples are taken with a sampling tube or fire shovel.

Coarse aggregate should be sampled in the following manner:

- 1) Locate the area of the stockpile from which plant loading will begin.
- 2) Using a front-end loader, dig into the stockpile and set aside a small pile of 10 to 15 t of material. This should be done in the same manner as the plant is being loaded. (Figs. 2-8 and 2-9) When forming the small pile, the loader bucket should be as low as possible and the operator should roll the material from the bucket rather than dumping the material. Reducing the distance the material is allowed to free-fall will reduce the amount of segregation that may occur in the small pile (Fig. 2-10). Each additional bucket load of material should be taken and dumped in the same manner as set out above and should be placed uniformly over the preceding one. (Fig. 2-11)

- Thoroughly mix the small pile. Using the loader bucket, go to the end of the oblong pile and roll the material over. Keeping the loader bucket as low as possible, push the bucket into the material until the front of the bucket passes the midpoint of the original pile. The loader bucket should then be slowly raised and rolled forward thus producing a smooth mixing of the material. (Figs. 2-12, 2-13, and 2-14) Go to the opposite end of the pile and repeat this mixing procedure. If the pile does not appear to be reasonably uniform additional mixing should be done.
- 4) The pile is now ready for sampling. Do <u>not</u> strike off the top. (Fig. 2-15) The sample is taken at the center of the volume which is approximately one-third of the height of the pile. The sample consists of not less than 6 full shovels of material taken at equal increments around the pile (Figs. 2-16, 2-17 and 2-18). The shovel is inserted full-depth horizontally into the material and raised vertically. Care should be taken to retain as much of the material as possible on the blade of the shovel. (Fig. 2-19)





Figure 2-9 Figure 2-9

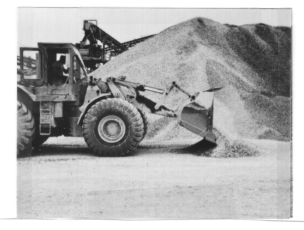




Figure 2-10 Figure 2-11





Figure 2-13 Figure 2-13





Figure 2-14









Figure 2-17



Figure 2-18



Figure 2-19

Fine aggregates are sampled using the same technique used to sample coarse aggregates, except a sampling tube or fire shovel is used instead of the large shovel.

Truck, Car, or Barge Sampling

Direct sampling from trucks, railroads cars, or barges is <u>not</u> recommended. There are a number of factors which may influence the quality of the material before it is used. Therefore, material being shipped by trucks, railroad cars, or barges should be sampled at the point of delivery after it has been stockpiled.

Size of Original Sample

The following is a list of recommended minimum sample sizes to be used as a guide when collecting materials.

MATERIAL	SAMPLE SIZE
No. 2 coarse aggregate	220 lb
No. 5 coarse aggregate	110 lb
No. 8 coarse aggregate	55 lb
No. 9 coarse aggregate	35 lb
No. 11 & No. 12 coarse aggregate	25 lb
All sands	25 lb

The weight of the sample depends on the maximum particle size of the material being tested. As a rule, a larger top size material requires a larger sample. A 35 lb sample of No. 5 coarse aggregate would not be as representative of that material as a 35 lb sample of natural sand. The size of sample for materials that do not meet a specific INDOT size should be the same as the INDOT size it's gradation is comparable to.

Reducing a Sample to Test Size

The total sample must be reduced to a sample size that can be quickly tested. Time will not allow the technician to run the total sample. The key to sample reduction is to ensure that the sample remains representative of the material in the stockpile. This practice is commonly referred to as splitting a sample. There are four different methods to reduce a sample to the proper test size.

1) Mechanical Splitter is the most accepted method of reducing to test size all coarse aggregate material smaller than gradation size No. 2;

- 2) Sand Splitter is the accepted method of fine aggregate that is drier than the saturated-surface-dry condition. As a quick check to determine this, if the sand retains its shape when molded in the hand, it is considered wetter than saturated-surface-dry;
- 3) Miniature Stockpile is the method used for fine aggregate that has free moisture on the particle surfaces; and
- 4) Quartering is the method that is used when a mechanical splitter is not available.

Mechanical Splitter

The Mechanical Splitter splits the sample into halves as the material passes through the spaces between the bars in the splitter. The same number of each particle size will go into each half of the sample, thus keeping the reduced sample representative of the total collected sample.

In using the Mechanical Splitter, adjust the splitter bars so that the bar opening is approximately twice the maximum particle size of the material to be split. A No. 5 aggregate has a maximum particle size of 1 1/2 in. Therefore, the recommended bar opening should be 3 in. or 6 bars wide (each bar is approximately 1/2 in.). INDOT allows the bar opening at 3 in. (6 bars) for all coarse aggregate No. 5 or smaller. The splitter must be level to ensure that each half of the split is approximately the same size; within approximately 10 percent of each other by weight.

The splitting procedure is as follows:

- 1) Properly place the pans under the splitter in such a way that all of the particles diverting in both directions will be caught;
- 2) Pour the sample evenly into the hopper;
- 3) Open the hopper fully and allow the material to free fall through the splitter;
- 4) If wet particles stick inside the splitter, gently tap the splitter with a rubber hammer to loosen them;
- 5) To ensure that the sample has not been segregated during sampling, place both halves of the sample back into the hopper and repeat the splitting operation; and

After the second splitting, the two receiving pans will contain approximately the same amount of material. Only one pan is placed back into the hopper and the splitting procedure repeated until a sample of the desired size is obtained. Skillful manipulation of the splitter will allow a sample of nearly any size to be made that is still representative of the material in the stockpile.

Sand Splitter

The sand splitter is a small version of the Mechanical Splitter except that the openings are fixed and there are no hopper doors.

The splitting procedure is as follows:

- 1) Place the pans under the splitter to catch all of the particles;
- 2) Slowly pour the dry sample into the splitter from the side (never from the end or corner);
- 3) Recombine the samples and split the sample a second time to eliminate any segregation; and
- 4) Reduce the sample to proper size by additional splitting of the material in one of the pans

Miniature Stockpile

This method is used for reducing all samples of fine aggregates when the material is in a damp or moist condition. If the sample to be split is dry, then the material must be moistened before using this method.

The splitting procedure is as follows:

- 1) Place the original sample on a clean, dry plate or other hard, smooth, non-absorptive surface;
- 2) Using a trowel or other suitable tool thoroughly mix the material to remove any segregation;
- 3) Shape the material into a conical pile; and
- 4) With a spoon or small trowel, randomly take at least five small portions of material around the pile and one-third way up the cone until the required test sample is obtained.

Quartering

Quartering is a non-mechanical method of reducing a sample. INDOT considers this the best method when a mechanical splitter is not available.

The quartering procedure is as follows:

- 1) Pour the sample in a conical pile in the center of a clean, dry, steel plate or other hard, smooth, non-absorptive surface;
- 2) Using a large trowel, shovel, or other suitable tool, thoroughly mix the material and reshape the sample into a conical pile;
- 3) Uniformly flatten the pile until the height is approximately equal to one-sixth the diameter;
- 4) With a large trowel or other suitable tool, divide the sample in half by vertically passing the tool through the center of the pile. In a similar manner divide each of these halves into two parts, thus quartering the sample; and
- 5) Combine diagonally opposite quarters of the material into two samples. Store one of these two halves. If the remaining material still weighs too much, repeat the entire quartering process until the proper test sample size is obtained.

Size of Test Sample (After Splitting)

The original sample must be reduced to test sample size which falls within the minimum and maximum weight in the following table.

AGGREGATE SIZE	MINIMUM (suggested)	MAXIMUM (suggested)
No. 2	11,300 g	
No. 5 and No. 8	6,000 g	8,000 g
No. 9	4,000 g	6,000 g
No. 11	2000 g	
No. 12	1,000g	
Fine Aggregate	300 g	

Gradation

Particle gradation is determined by a sieve or gradation analysis of aggregate samples. A sieve analysis involves passing the sample through a series of sieves, each of which has openings of specific sizes (Figure 2-20). Sieves are designated by the size of their openings. Coarse particles are trapped in the upper sieves; medium-sized particles pass through to the mid-level sieves; fines pass through to the lowest sieves.

The aggregate gradation considers the percentage by weight of the total sample that passes through each sieve. This is determined by weighing the contents of each sieve following the sieve analysis, then calculating the percentage passing each sieve.

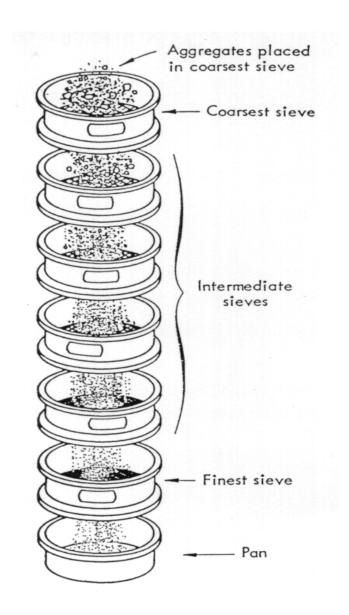


Figure 2-20. Sieve Analysis

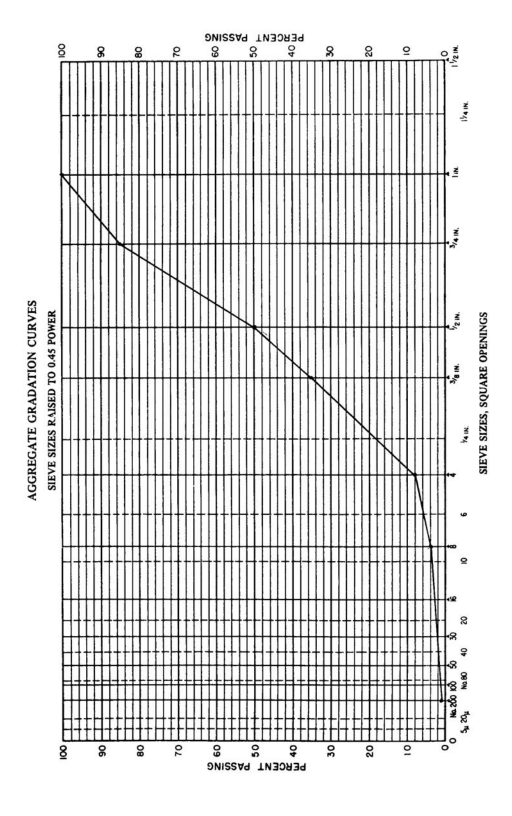


Figure 2-21. Gradation Chart

For the purpose of description, certain terms are used in referring to aggregate fractions. They are:

Coarse aggregate - Material that has a minimum of 20 percent retained on the No. 4 sieve.

Fine aggregate - Material that is 100 percent passing the 3/8 in. sieve and a minimum of 80 percent passing the No. 4 sieve.

Nominal Maximum Particle Size - Smallest sieve opening through which the entire amount of the aggregate is permitted to pass.

Maximum Particle Size - Largest sieve size listed in the specification through which all material must pass.

Aggregate gradation specifications for a given project can be presented graphically; Figure 2-21 is a typical gradation chart. On the chart, sieve sizes are presented horizontally and percent passing each sieve is shown vertically.

Sieve Analysis Test

Sieve analysis is used primarily to determine the particle-size distribution of materials. The results determine compliance of the particle-size distribution with the applicable requirements. The test is conducted on both the fine and coarse aggregates and is performed according to **AASHTO T 27**, with exceptions as noted in Section **904.06** of the Standard Specifications.

The procedure for a sieve analysis is as follows:

- 1) The dried sample is placed in the top sieve of properly nested sieves. The sieves are nested in sequence with the smallest sieve placed on the pan and stacked by increasing size.
- 2) The shaking time must be sufficient to ensure that the sample is divided into fractional sizes. The actual shaking time is required to be determined in accordance with **ITM 906**. The following times are minimum for shakers used by the industry.

Coarse Aggregate, Size 9 or larger 5 Minutes Coarse Aggregate, Smaller than Size 9 10 Minutes Fine Aggregates 15 Minutes 3) At the conclusion of sieving, the material retained on each sieve is carefully transferred to a weigh pan and weighed. The weight retained of the material on each sieve is recorded on the Gradation Analysis sheet. The weight cannot exceed the allowable amount on each sieve as indicated in Table 1.

Note: The larger sieves (above the No. 16) are cleaned with a small trowel or piece of flat metal. The sieves between the No. 16 and No. 50 are cleaned with a wire brush. Sieves under the No. 50 are cleaned with a soft bristle brush. Care must be taken not to damage the sieves.

4) The weight passing each sieve is calculated next by subtracting the weight retained on the largest sieve from the total sample weight. The weight retained on the next largest sieve is subtracted from the weight of material still remaining from the first subtraction. This process is continued for all sieves.

Example:

1 in.	5942.1 - 690.6	=	5251.5
3/4 in.	5251.5 - 2492.7	=	2758.8
3/8 in.	2758.8 - 1397.1	=	1361.7
No. 4	1361.7 - 997.0	=	364.7
No. 8	364.7 - 264.5	=	100.2
Pan material		=	88.1

TABLE 1 APPROXIMATED SIEVE OVERLOAD

SCREEN SIZE	STANDARD 15 in. x 23in.	STANDARD 14 in. x 14 in.	12 in. DIAMETER	8 in. DIAMETER
3 in.	40.5 kg	23.0 kg	12.6 kg	
2 in.	27.0 kg	15.3 kg	8.4 kg	3.6 kg
1-1/2 in.	20.2 kg	11.5 kg	6.3 kg	2.7 kg
1 in.	13.5 kg	7.7 kg	4.2 kg	1.8 kg
3/4 in.	10.2 kg	5.8 kg	3.2 kg	1.4 kg
1/2 in.	6.7 kg	3.8 kg	2.1 kg	890 g
3/8 in.	5.1 kg	2.9 kg	1.6 kg	670 g
No. 4	2.6 kg	1.5 kg	800 g	330 g

8 in. diameter sieves, No. 8 to No. 200 shall not exceed 200g / sieve

12 in. diameter sieves, No. 8 to No. 200 shall not exceed 469g / sieve

TOTAL WE	IGHT 5	5942.1 g				
SIEVE SIZE		GRAD. HT RET.	WEIGHT RETAINED	WEIGHT PASSING	PERCENT PASSING	PERCENT REQUIRED
2 1/2 in.			g	g	%	%
2 in.			g	g	%	%
1 1/2 in.			g	g	%	%
1 in.			0 g	5942.1 g	%	%
3/4 in.			690.6 g	5251.5 g	%	%
1/2 in.		Т	2492.7 g	2758.8 g	%	%
3/8 in.	P F		1397.1 g	1361.7 g	%	%
No. 4		g	997.0 g	364.7 g	%	%
No. 6		g	g	g	%	%
No. 8		g	264.5 g	100.2 g	%	%
No. 16		g	g	g	%	%
No. 30		g	g	g	%	%
No. 50		g	g	g	%	%
No. 100		g	g	g	%	%
No. 200		g	g	g	%	%
PAN		g	88.1 g	g	%	%
DECANT	ORIC	GINAL	FINAL	GRAMS LOSS	PERCENT LOSS	PERCENT REQUIRED
		g	g	g	%	%

5) The percent passing is calculated for each sieve by using the following formula:

% Passing =
$$\frac{\text{Weight passing each sieve}}{\text{Original dry sample weight}} \times 100$$

Example:

$$\frac{5251.5}{5942.1}$$
 x 100 = 88.4%
1/2 in. $\frac{2758.8}{5942.1}$ x 100 = 46.4% etc.

6) If the test has been done accurately, the sum of all the fractional weights retained (including the material in the pan) will be approximately equal to the original dry weight. If the two weights differ by more than 0.3 percent, based on the original dry sample weight, the results are considered invalid.

Original Dry Weight - Summation Weights Measured x100
Original Dry Weight

Example:

$$\frac{5942.1 - (690.6 + 2492.7 + 1397.1 + 997.0 + 264.5 + 88.1)}{5942.1} \times 100 =$$

0.2% = valid test

NOTE: This is a check on workmanship and does not have to be performed every time. Occasionally perform this calculation to be assured that the proper testing technique is being used.

TOTAL WE	IGHTS	5942.1 g				
SIEVE SIZE		GRAD. HT RET.	WEIGHT RETAINED	WEIGHT PASSING	PERCENT PASSING	PERCENT REQUIRED
2 1/2 in.	-		g	g	0/0	9/0
2 in.	-		g	g	%	9/0
1 1/2 in.	-		g	g	%	9/
1 in.	-		0 g	5942.1 g	100 %	9/
3/4 in.	-		690.6 g	5251.5 g	88.4%	0/
1/2 in.		T	2492.7 g	2758.8 g	46.4%	0/
3/8 in.	P F		1397.1 g	1361.7 g	22.9%	0/
No. 4		g	997.0 g	364.7 g	6.1%	9/
No. 6		g	g	g	%	9/
No. 8		g	264.5 g	100.2 g	1.7%	9/
No. 16		g	g	g	%	9/
No. 30		g	g	g	%	0/
No. 50		g	g	g	%	9/
No. 100		g	g	g	%	9/
No. 200		g	g	g	%	0/
PAN		g	88.1 g	g	%	0/
DECANT	ORIC	GINAL	FINAL	GRAMS LOSS	PERCENT LOSS	PERCENT REQUIRED
		g	g	g	%	9/

Moisture Test

When aggregates are obtained from the cold feed belt or the belt discharge for a drum mix HMA plant, the moisture of the aggregates must be determined to adjust aggregate weights for moisture content and to determine the amount of binder to add to the HMA.

For moisture content the sample must be reduced to test size and the test run as quickly as possible after the sample has been taken. Any delay in running the test after the sample has been obtained may allow the material to lose moisture and cause inaccurate results.

The test procedure is:

- 1) Weigh the sample before drying and record the weight;
- 2) Dry the sample and allow to cool to room temperature;
- 3) Weigh the sample and record the weight; and
- 4) Determine the moisture percent using this formula:

% Moisture =
$$\frac{\text{Weight Wet - Weight Dry}}{\text{Weight Dry}} \times 100$$

Certified Aggregate Producer Program

The Certified Aggregate Producer Program (CAPP) is a program in which a qualified mineral aggregate Producer desiring to supply material for INDOT use may do so by assuming all of the Plant site controls and a portion of the testing responsibility that had been previously assumed by INDOT.

The benefit of the CAPP is that the Producer is providing material that has a consistent gradation. Documentation of gradation test results for both during production and as the aggregate is loaded-out is available at any time. This information can save the Producer time and money in designing HMA. Also, problems that may occur during production of the HMA due to inconsistent gradations may be quickly traced with information readily available at the aggregate source on all aspects of production of the material.

ASPHALT

Asphalt is a black, cementing material that varies widely in consistency from solid to semisolid (soft solid) at normal air temperatures. When heated sufficiently, asphalt softens and becomes a liquid, which allows it to coat aggregate particles during HMA production.

Asphalt is made up largely of a hydrocarbon called bitumen. Consequently, asphalt is often called a bituminous material. Virtually all asphalt used in the United States is produced by modern petroleum refineries and is called petroleum asphalt. The degree of control allowed by modern refinery equipment permits the production of asphalts with specific characteristics suited to specific applications. As a result, different asphalts are produced for paving, roofing and other special uses.

Paving asphalt, commonly called binder, is a highly viscous (thick), sticky material. It adheres readily to aggregate particles and is therefore an excellent cement for binding together aggregate particles in a HMA. The binder is an excellent waterproofing material and is unaffected by most acids, alkalies (bases) and salts. This means that a properly constructed HMA pavement is waterproof and resistant to many types of chemical damage.

Binder for paving can also contain modifiers to improve performance properties. Some of these binders require special storage and handling. The material suppliers recommendations should be followed to insure that these performance characteristics are not altered or lost before mixing and placement of the HMA.

Binder changes when it is heated and/or aged. It tends to become hard and brittle and therefore lose some of its ability to adhere to aggregate particles. These changes can be minimized by understanding the properties of the binder and taking steps during construction to ensure that the finished pavement is built in a way that will retard the aging process.

Source and Nature of Asphalt

Because asphalt is used for many purposes, there is sometimes confusion about where asphalt comes from, how it is refined, and how it is classified into grades. There is similar confusion about terms related to asphalt properties and use. It is the purpose of this section to discuss the sources and nature of paving asphalt in sufficient detail for a clear understanding of fundamental concepts.

Petroleum Refining

Crude petroleum is refined by distillation, a process in which various fractions (products) are separated out of the crude. Distillation is accomplished by raising the temperature of the crude petroleum in stages. As shown in Figure 2-22, different fractions separate at different temperatures.

The lighter fractions are separated by simple distillation. The heavier distillates, often referred to as gas oils, can be separated only by a combination of heating and applying a vacuum. The product which cannot be distilled under vacuum distillation is asphalt.

Figure 2-23 is a schematic illustration of a typical refinery. It shows the flow of petroleum during the refining process.

Asphalt Refining

Different types of asphalt are required for different applications. To produce asphalts that meet specific requirements, refiners must have a way to control the properties of the asphalts they produce. This is often accomplished by blending crude petroleums of various types together before processing. Blending allows refiners to combine crudes that contain asphalts of varying characteristics in such a way that the final product will have exactly the characteristics required by the asphalt user.

Once the crude petroleums have been blended together, there are two widely used processes by which asphalt can be produced from them: vacuum distillation and solvent extractions.

As discussed above, vacuum distillation involves separating the asphalt from the crude by applying heat and a vacuum. In the solvent extraction process, additional gas oils are removed from the crude, leaving residual asphalt.

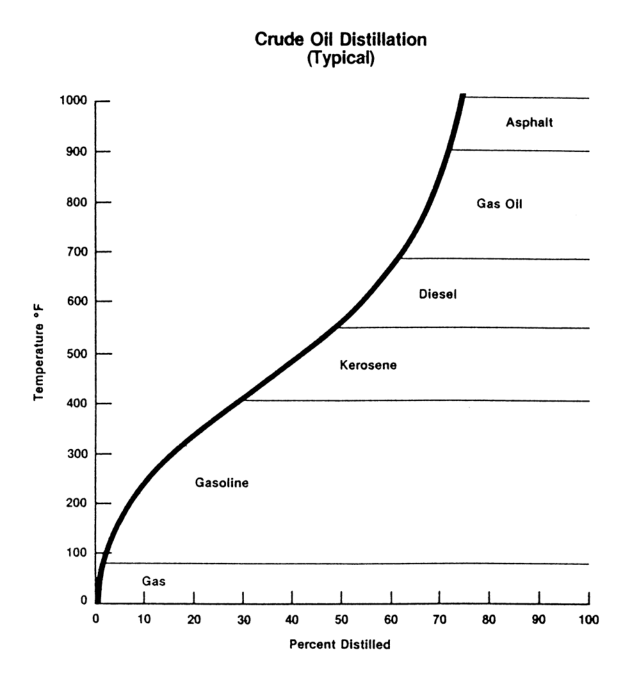


Figure 2-22. Typical Distillation Temperatures and Products

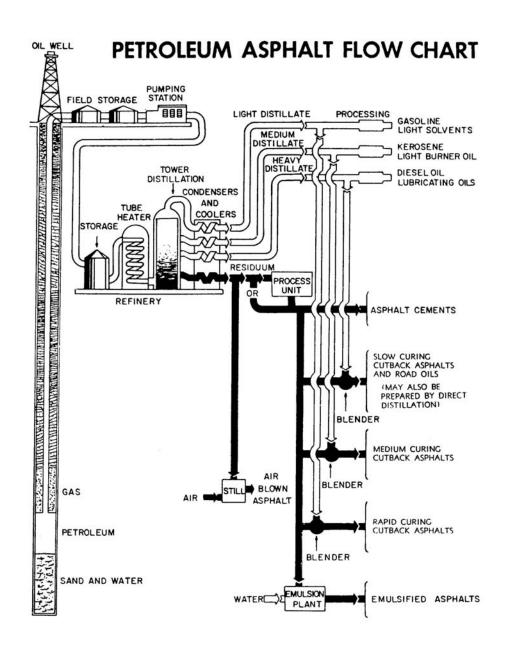


Figure 2-23. Typical Refining Process

2-34

Performance Graded Binder

In 1987, the Strategic Highway Research Program (SHRP) began developing a new system for specifying asphalt materials. The final product of the SHRP asphalt research program is a system called Superpave, short for Superior Performing Asphalt Pavements. Superpave represents an improved system for specifying binder and mineral aggregates, developing HMA designs, and analyzing and establishing pavement performance prediction. The system includes a binder specification, a HMA design and analysis system, and computer software that integrates the system components. The unique feature of the Superpave system is that it is a performance-based specification system. The tests and analyses have direct relationships to field performance.

The SHRP binder specifications are based on fundamental properties which are measured at actual in service temperatures where the critical distresses occur. The upper temperature extreme is designated as the average 7-day maximum pavement design temperature. This temperature is obtained by accumulating the temperature from each successive 7-day period throughout the summer, and choosing the 7-day period which yields the largest average. The lower temperature extreme is designated as the minimum pavement design temperature. The temperatures for the specification were determined by gathering data from more than 1,000 weather stations and are available with the SHRP mix design method, SUPERPAVE. An example of how this grading system works is shown in Figure 2-24. The specification limits for the PG Binders are listed in 902.01 of the Standard Specifications.

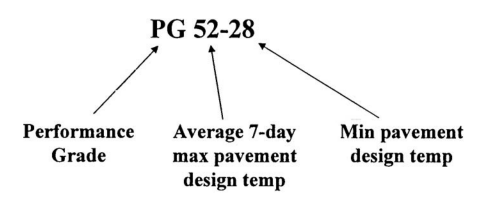


Figure 2-24. PG Grading System

The PG asphalt binder specification is based on the following tests:

- 1. Aging
- 2. Dynamic Shear
- 3. Direct Tension
- 4. Bending Beam
- 5. Viscosity
- 6. Flash Point

Asphalt Binder Aging

Aging evaluation of the binder is performed by both the Rolling Thin Film Oven (RTFO) (Figure 2-25) and Pressure Aging Vessel (PAV) (Figure 2-26). The RTFO test is to be performed in accordance with **AASHTO T 240** to simulate aging that occurs during construction.

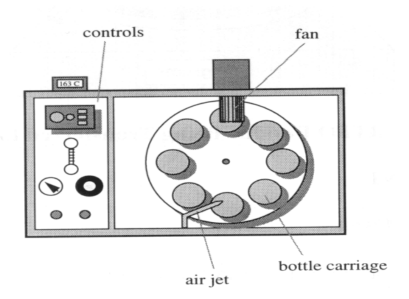


Figure 2-25. Rolling Thin Film Oven (RTFO)

The PAV test (AASHTO R 28) is designed to simulate long-term in-service aging (8-10 years), and thereby provide a procedure to estimate the binder properties after aging. The PAV uses a pressurized container into which pans containing RTFO residue binders are placed. Samples in the PAV are subjected for 20 hours to a specified pressure and to variable temperatures depending on the climate of the region.

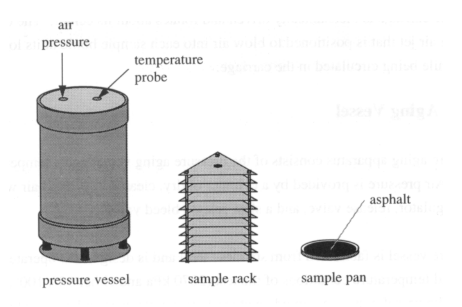


Figure 2-26. Pressure Aging Vessel

Dynamic Shear

The dynamic shear test (**AASHTO T 315**) provides data on the visco-elastic or rheological properties of the binder. Rheology is defined as "the study of deformation and flow of matter", in this case binder. The Dynamic Shear Rheometer (DSR) is used to evaluate binders at upper and intermediate service temperature ranges, where rutting and fatigue are the respective primary distress mechanisms. Although rutting and fatigue cracking properties of HMA are not controlled solely by the binder, the information generated from this test is valuable in gaining a better understanding of the visco-elastic properties of the binder. Figure 2-27 gives a schematic view of how the DSR functions.

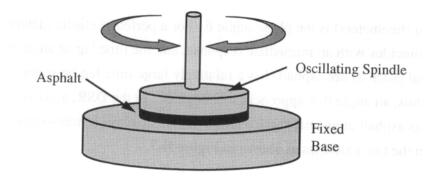


Figure 2-27. Dynamic Shear Rheometer

Direct Tension

The direct tension test (**AASHTO T 314**) pulls a "dogbone" shaped specimen of binder to failure at a controlled rate of "stretching" or strain. The maximum load to failure (failure stress) as well as the maximum amount of "stretch" to failure (failure strain) are recorded. The test is used to estimate low temperature failure properties of the binder. A drawing of the test specimen is shown in Figure 2-28.

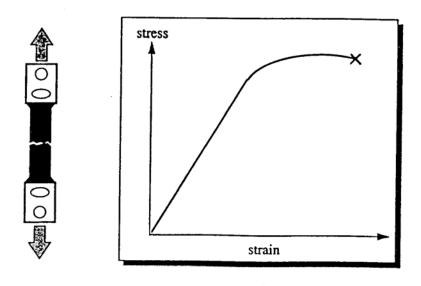


Figure 2-28. Direct Tension

Bending Beam

Low temperature stiffness of the binder is evaluated using the bending beam rheometer. A constant load is applied to a beam sample of the binder which is submerged in a bath. Deflections of the beam are measured and a stiffness versus time curve is developed.

Data generated from this test (AASHTO T 313) relates to low temperature thermal cracking. Figure 2-29 provides a schematic view of the test apparatus.

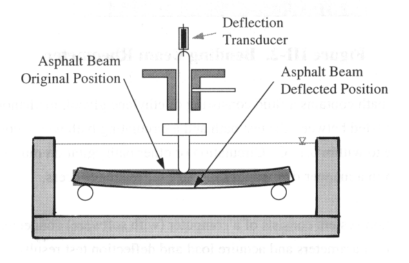


Figure 2-29. Bending Beam

Viscosity

The rotational viscosity test (**AASHTO T 316**) consists of turning a spindle in a sample of binder and reading viscosity as the resistance to movement of the spindle. The test is performed at high temperatures to ensure proper handling of the binder. A low temperature test can also be performed to develop a viscosity-temperature curve. This curve can then be used to develop mixing and compaction temperatures for the specific binder. Figure 2-30 illustrates the test apparatus.

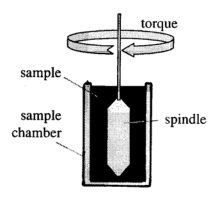


Figure 2-30. Brookfield Viscometer

Flashpoint

The flashpoint of a binder is the lowest temperature at which volatile materials separate from a sample in sufficient concentration to "flash" in the presence of an open flame. Flashpoint must not be confused with firepoint, which is the lowest temperature at which the binder will catch fire and burn. Flashpoint involves only instantaneous combustion of the volatile fractions separating from the binder.

The flashpoint of a binder is determined to identify the maximum temperature at which it can be handled and stored without danger of flashing. This is important information since binder is usually heated in storage to keep its viscosity low enough so that the material can be pumped.

The procedure for determining flashpoint is to gradually heat a sample of binder in a brass cup while periodically holding a small flame over the surface of the sample (Figure 2-31). The temperature at which an instantaneous flashing of vapors occurs across the surface is taken to be the flashpoint. The Cleveland Open Cup Test (AASHTO T 48) is the procedure for determining the flashpoint.

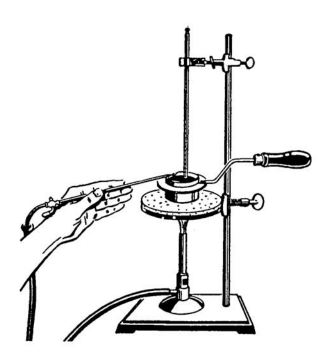


Figure 2-31. Flashpoint Test; Cleveland Open Cup Test

ASPHALT HANDLING, STORAGE, AND SAMPLING

The safety record for handling, storing and sampling asphalt materials is good. Nonetheless, there have been accidents resulting in property damage, personal injury and loss of life. To prevent such mishaps, everyone must know and follow good safety practices. When an accident does occur, everyone must know how to react and what first aid treatment is appropriate.

The technician should also be aware of the potential sources of contamination that might exist where asphalt materials are stored or handled. Changes in volume that the asphalt material undergoes when heated or cooled should also be understood. This knowledge is especially important when comparing asphalt material quantities measured at different temperatures.

Handling

At a HMA plant, asphalt material temperatures may exceed 300°F. Metal surfaces of plant equipment often range between 150°F and 200°F. Consequently, momentary contact with hot asphalt materials or with plant equipment, including tanks, pipelines, dryers, boilers, and boiler houses, can cause severe burns to exposed flesh. Four general precautions against such burns are:

- 1) Be aware of where burn hazards are located;
- 2) Use designated walk areas and stay clear of hazardous situations;
- 3) Always wear appropriate work clothing.; and
- 4) Know and follow all plant safety procedures related to handling hot material and equipment.

If a burn does occur, follow these general treatment guidelines:

- 1) In the case of local asphalt material skin burns, apply cold water or an ice pack to reduce the heat in the asphalt material and the skin;
- In cases where burns cover more than 10 percent of the body (approximately the skin area of one arm or half a leg), apply lukewarm water instead of cold water. Lukewarm water will reduce the temperature of the asphalt material and skin without causing shock, which could be induced by applying cold water or ice to major burns;
- 3) Do not remove the asphalt material from the skin;

FIRST AID FOR MOLTEN ASPHALT **CEMENT BURNS**

In the event of a MOLTEN ASPHALT CEMENT BURN:

COOL the asphalt cement and affected parts of the body immediately.

Methods of cooling (in order of preference):

- 1. Completely submerge affected area in ice water,
- 2. Completely submerge affected area in tap water,
- 3. Place affected area under running water.

DO NOT DELAY

Use any available water, cooler than body temperature, while arranging for better cooling.

CAUTION: DO NOT apply Ice directly to affected area.

LEAVE cooled asphalt cement on affected area.

Proceed with the following:

MINOR ASPHALT CEMENT BURNS-at first opportunity get victim to physician.

injury to small areas of fairly insensitive flesh Involving a small quantity of asphalt cement.



Hospital_ Clinic Physician's Office.

injury to the head, face or extremitles;

Injury when large amounts of asphalt cement are involved; Evidence of nausea or faintness.

TREATMENT FOR SHOCK

In the event shock occurs, do the following:

- 1. Keep victim lying down and quiet.
- 2. Keep victim covered with a blanket or something similar to keep body temperature at normal, 98°F (37°C).
- 3. Keep victim's head lower than feet to promote blood supply to head and chest.

DO NOT ATTEMPT TO REMOVE THE ASPHALT CEMENT with products containing solvents or ammonia. Natural separation will occur in about 48-72 hours.

If necessary, for early removal, soak bandage in mineral oil and place over affected area for 2 to 3 hours.

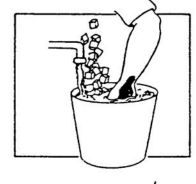




Figure 2-32

- 4) Do not cover the burned area with a bandage; and
- 5) Have a physician examine the burn immediately.

Hydrogen sulfide is a product of the reaction between hydrogen and sulphur naturally present in asphalt materials. In low concentration, hydrogen sulfide is not dangerous; however, in the high concentrations sometimes found in storage tanks and other closed areas, it can be lethal. To prevent overexposure to hydrogen sulfide fumes, follow these guidelines:

- 1) Keep your face at least two feet away from asphalt material tank hatch openings;
- 2) Stay upwind of open hatches; and
- 3) Avoid breathing fumes when opening hatch covers or taking samples.

In case of overexposure to hydrogen sulfide fumes, do the following:

- 1) Move victim immediately to fresh air;
- 2) Administer oxygen if breathing is difficult;
- 3) Start artificial respiration if breathing stops; and
- 4) Have the victim examined by a physician immediately.

See Figure 2-32 for first aid information.

Storage

The Certified Technician is not required to test the binder or asphalt emulsion for specification purposes. However, the technician should be aware that these materials are tested for compliance from the HMA plant and not the asphalt refinery. It is therefore important that the technician know what plant conditions can change the physical properties of these materials and possibly cause the materials to become out of specifications.

Binder

Three major problems can cause the asphalt binder to shift outside of specification. The first is contamination. If hot oil is used to heat the binder tank then any leak in the system can contaminate the binder. The contamination results in a lower viscosity binder. Another source of contamination is fuel oil. If binder is contaminated with significant quantities of fuel oil because of flushing pumps and lines then the binder stiffness drops.

A second and more prevalent problem is the effect that extended heating in storage tanks has on the binder. When binder is heated, it slowly increases in stiffness. The rate of change in viscosity is dependent on the size of tank, surface area, amount of circulation, temperature, etc. The problem of compliance generally is found after long rainy periods with extended heating. The failures can be minimized by reducing heating and circulation during these periods of non-production. When the technician has some doubts about the material, the binder should be tested to determine compliance. This will greatly minimize the number of problems and penalties associated with failures.

A third problem encountered with binder is improper storage. Modified binders include additives to improve (extend) the grade to encompass a wider temperature range for performance. Many of these modified binders must be stored under special conditions to maintain their properties. Material supplier recommendations for storage and handling should be followed.

Sampling

Normally binder samples are taken from a sampling valve on a storage tank. Here are a few important rules to follow when sampling the binder.

- 1) To ensure that samples are representative of the entire shipment, take them from the sampling valves provided for that purpose (Figure 2-33). Dip samples taken from the top of a tank are not normally representative. Other sampling methods are described in **AASHTO T 40**;
- 2) Use only new, clean dry sample containers;
- 3) Allow at least 4 quarts of the binder to drain out of the valve before taking samples. This cleans out the valve and the lines and helps to provide a representative sample;
- 4) Seal filled containers immediately with clean, dry, tight fitting lids. Wipe any spilled material from the container with a clean, dry cloth; NEVER with a cloth dipped or soaked in solvent;
- 5) Label all containers clearly. Do not label container lids, because once the labeled lid is removed, it will be difficult to identify the sample in the container. Do all labeling with a wick marking pen. Use tags only when there is no danger of their being lost in transit; and
- 6) Follow all safety precautions for handling and storing hot binder. Remember, the binder is hot when sampled, so wear protective clothing (gloves, face shield, long-sleeved shirt) to protect from burns and splattering.

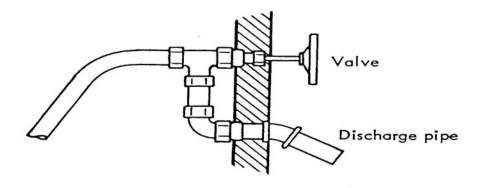


Figure 2-33. Sampling Device for Storage Tanks

BINDER VOLUME

Specific Gravity

Specific gravity is the ratio of the weight of a volume of material to the weight of an equal volume of water, both at a specified temperature. As an example, a substance with a specific gravity of 1.6 weighs 1.6 times as much as an equal volume of water.

Knowing the specific gravity of the binder being used is important for two reasons. First, binder expands when heated and contracts when cold. This means that the volume of a given amount of binder will be greater at higher temperatures than at lower ones. Specific gravity measurements provide a yardstick for making temperature-volume corrections, which is discussed later.

Second, specific gravity of the binder is essential in the determination of the percentage of voids (air spaces) in the compacted pavement. (Chapter 4).

Specific gravity is usually determined by the pycnometer method as shown in Figure 2-34. Because specific gravity varies with the expansion and contraction of binder at different temperatures, results are normally expressed in terms of Specific Gravity at a given temperature for both the material and the water used in the test. (Example: Sp.Gr. 1.05 at 60°F/60°F means that the specific gravity of the binder is 1.05 when both the binder and the water are at 60°F).

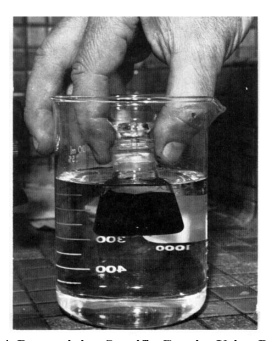


Figure 2-34. Determining Specific Gravity Using Pycnometer

Temperature Volume Relationships

Changes in volume of the binder can be a source for confusion in the plant when binder must be converted from weight to volume. When binder is shipped and stored, the basis of payment is weight. This eliminates confusion in payment. However, binder in some plants is metered rather than weighed. When this occurs the technician must know how to convert weight to volume and vice versa. The density or specific gravity of binder is the ratio of weight to volume at a specified temperature. Normally this temperature is 60°F. As the temperature of the binder increases the weight to volume ratio drops. It is important to understand in the plant, if the binder is being metered, the density of the binder at 60°F and the temperature of the binder. It is also important to know if the meter is compensating for this temperature and converting the registered volume back to 60°F. If not, then the binder content in the HMA will be less than expected.

Another use of temperature volume relationships is to determine inventory. If the storage tank is gauged and a volume is determined then that volume must be reduced to a volume at 60°F. This value can then be converted to weight easily. Examples of converting a tank gauge volume to volume at 60°F and then to weight are as follows:

Example

Volume of binder is measured to be 6505 gallons at a temperature of 290°F. The specific gravity of the binder is 1.021. Determine the volume of binder at the standard base temperature of 60°F.

Volume correction factor of 0.9220 is read for the observed temperature of 290°F.

Vol. $@60^{\circ}F = Vol. @(actual temp.) \times Vol. Conversion Factor$

Volume @
$$60^{\circ}$$
F = 6505×0.9220
= 5998 gallons

To convert volume at 60°F to Tonnage:

Tonnage =
$$\underline{\text{Vol.}}$$
 @ 60°F x Specific Gravity @ 60°F x 8.33 lb/gal 2000 lb/t

Tonnage =
$$\underline{5998 \times 1.021} \times 8.33 = 25.5 \text{ t}$$

 $\underline{2000}$

GROUP O (°F)

GROUP 0-SPECIFIC GRAVITY AT 60°F ABOVE 0.966

LEGEND: t = observed temperature in degrees Fahrenheit
M = multiplier for correcting oil volumes to the basis of 60°F

	M	1		1			nes to the	$\overline{}$	
-		-	M	1	M		M	1	M
0	1.0211	50	1.0035	100	0.9861 0.9857	150	0.9689	200	0.9520 0.9516
2	1.0204	52	1.0028	102	0.9854	152		202	0.9513
3	1.0201	53	1.0024	103	0.9851	153	0.9679	203	0.9509
5	1.0197	54	1.0021	104	0.9847	154 155	0.9675	204	0.9506
6	1.0190	56	1.0017	106	0.9840	156	0.9672	205	0.9503
7	1.0186	57	1.0010	107	0.9837	157	0.9665	207	0.9496
8	1.0183	58	1.0007	108	0.9833	158 159	0.9662 0.9658	208	0.9493
10	1.0176	60	1.0000	110	0.9826	160	0.9655	210	0.9486
11	1.0172	61	0.9997	111	0.9823	161	0.9652	211	0.9483
12	1.0169	62	0.9993	112	0.9819	162 163	0.9648	212	0.9479 0.9476
14	1.0162	64	0.9986	114	0.9813	164	0.9641	214	0.9472
15	1.0158	65	0.9983	115	0.9809	165	0.9638	215	0.9469
16	1.0155	66	0.9979	116	0.9806	166	0.9635	216	0.9466
17	1.0151	67 68	0.9976 0.9972	117	0.9802 0.9799	167	0.9631	217	0.9462
19	1.0144	69	0.9969	119	0.9795	169	0.9624		0.9456
20	1.0141	70	0.9965	120	0.9792	170	0.9621	220	0.9452
21 22	1.0137	71 72	0.9962	121	0.9788	171 172	0.9618	221	0.9449
23	1.0130	73	0.9955	123	0.9782	173	0.9611	223	0.9442
24	1.0126	74	0.9951	124	0.9778	174	0.9607	224	0.9439
25 26	1.0123	75 76	0.9948	125 126	0.9775 0.9771	175 176	0.9604	225 226	0.9436
27	1.0116	77	0.9941	127	0.9768	177	0.9597	227	0.9429
28	1.0112	78	0.9937	128	0.9764	178	0.9594	228	0.9426
29 30	1.0109	79 80	0.9934	129	0.9761	179	0.9590	229	0.9422
31	1.0102	81	0.9927	130 131	0.9758	180 181	0.9587	230	0.9419
32	1.0098	82	0.9923	132	0.9751	182	0.9580	232	0.9412
33 34	1.0095	83 84	0.9920	133 134	0.9747	183 184	0.9577	233	0.9409
35	1.0088	85	0.9913	135	0.9740	185	0.9570	235	0.9402
36	1.0084	86	0.9909	136	0.9737	186	0.9567	236	0.9399
37 38	1.0081	87 88	0.9906	137 138	0.9734	187	0.9563	237	0.9395
39	1.0074	89	0.9899	139	0.9727	188 189	0.9560 0.9557	238 239	0.9392
40	1.0070	90	0.9896	140	0.9723	190	0.9553	240	0.9385
41	1.0067	91 92	0.9892	141	0.9720	191	0.9550	241	0.9382
43	1.0060	93	0.9885	142	0.9713	192 193	0.9547	243	0.9379
44	1.0056	94	0.9882	144	0.9710	194	0.9540	244	0.9372
45	1.0053	95	0.9878	145	0.9706	195	0.9536	245	0.9369
46 47	1.0049	96 97	0.9875	146	0.9703	196 197	0.9533		0.9365 0.9362
48	1.0042	98	0.9868	148	0.9696	198	0.9526	248	0.9359
49	1.0038	99	0.9864	149	0.9693	199	0.9523	249	0.9356

Figure 2-35. Temperature Volume Corrections for Binder

GROUP O continued (OF)

GROUP O-SPECIFIC GRAVITY AT 60°F ABOVE 0.966

LEGEND: t = observed temperature in degrees Fahrenheit
M = multiplier for correcting oil volumes to the basis of 60°F

-	M -	1	M	1	M	7			
250		-		-		1	M	1	M
251		300	0.9187	350	0.9024	400	0.8864	450	0.8705 0.8702
252		302	0.9181	352	0.9018	402	0.8857	452	0.8699
253 254		303	0.9177	353 354		403	0.8854	453	0.8696
255		305	0.9171	355	0.9008	404	0.8851	454	0.8693
256	0.9332	306	0.9167	356	0.9005	406	0.8845	456	0.8690 0.8687
257 258		307	0.9164	357	0.9002	407	0.8841	457	0.8683
259		309	0.9161	358 359	0.8998	408	0.8838	458	0.8680
260	0.9319	310	0.9154	360	0.8992	410	0.8832	460	0.8674
261	0.9316	311	0.9151	361	0.8989	411	0.8829	461	0.8671
262 263		312	0.9148	362 363	0.8986	412	0.8826 0.8822	462	0.8668
264		314	0.9141	364	0.8979	414	0.8819	464	0.8661
265		315	0.9138	365	0.8976	415	0.8816	465	0.8658
266 267	0.9299	316	0.9135	366	0.8973	416	0.8813	466	0.8655
268	0.9293	318	0.9128	368	0.8966	418	0.8806	468	0.8649
269	0.9289	319	0.9125	369	0.8963	419	0.8803	469	0.8646
270 271	0.9286	320	0.9122	370	0.8960	420	0.8800	470 471	0.8643
272	0.9279	322	0.9115	372	0.8953	422	0.8794	472	0.8640
273	0.9276	323	0.9112	373	0.8950	423	0.8791	473	0.8633
274 275	0.9273	324	0.9109	374	0.8947	424	0.8787	474	0.8630
276	0.9266	326	0.9103	375 376	0.8944	425	0.8784	475 476	0.8627 0.8624
277	0.9263	327	0.9099	377	0.8937	427	0.8778	477	0.8621
278	0.9259	328	0.9096	378 379	0.8934	428	0.8775 0.8772	478	0.8618
280	0.9253	330	0.9089	380	0.8928	430	0.8768	480	0.8615
281	0.9250	331	0.9086	381	0.8924	431	0.8765	481	0.8608
282 283	0.9246	332	0.9083	382	0.8921	432	0.8762	482	0.8605
284	0.9240	334	0.9076	384	0.8915	434	0.8759 0.8756	483 484	0.8602 0.8599
285	0.9236	335	0.9073	385	0.8912	435	0.8753	485	0.8596
286 287	0.9233	336 337	0.9070	386 387	0.8908	436 437	0.8749	486	0.8593
288	0.9227	338	0.9063	388	0.8902	438	0.8746	0.000	0.8590 0.8587
289	0.9223	339	0.9060	389	0.8899	439	0.8740		0.8583
290 291	0.9220	340	0.9057	390 391	0.8896	440	0.8737		0.8580
292	0.9213	342	0.9050	392	0.8892	441	0.8734		0.8577 0.8574
293	0.9210	343	0.9047	393	0.8886	443	0.8727	493	0.8571
294 295	0.9207	344	0.9044	394	0.8883	444	0.8724		0.8568
296	0.9200	346	0.9040	395 396	0.8880	445	0.8721		0.8565 0.8562
297	0.9197	347	0.9034	397	0.8873	447	0.8715	497	0.8559
298 299	0.9194	348 349	0.9031	398 399	0.8870	448	0.8712		0.8556
	5.7.70	347	0.7020	377	0.6607	747	0.8709	499	0.8552

Temperature Volume Corrections for Binder (continued)

Asphalt Supplier Certification Program

The Asphalt Supplier Certification Program (ASC) is a program in which a qualified asphalt supplier is permitted to manufacture and ship PG binders without complete pre-testing of the PG binder by INDOT or the supplier. PG binders are required to be supplied by an approved supplier in accordance with **ITM 581**.

PG binders that are shipped to HMA plants are required to have a shipping report indicating that the material was manufactured in accordance with the ASC program. Also, the supplier is required to furnish instructions with each PG binder concerning the proper storage and handling of the material.